STEVEN M. FOLTZ

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EDUCATION

PhD, Washington University in St. Louis, St. Louis, MO

June 2014 – present

Division of Biology and Biomedical Sciences (DBBS)

Expected defense: Winter 2019

Human and Statistical Genetics Program

Precision Medicine Pathway

Advisor: Li Ding, PhD

Thesis: Multi-platform genomics and transcriptomics integration for gene fusion discovery and

somatic mutation haplotyping in cancer

MS Biostatistics, University of Washington, Seattle, WA September 2011 – August 2013

Advisors: Brian Browning, PhD, and Sharon Browning, PhD

Thesis: Rare variant method for identity-by-descent detection in sequence data, and the time to most recent common ancestor given identity-by-descent segment length

BS Mathematics, Presbyterian College, Clinton, SC

August 2006 – April 2010

Advisor: Doug Daniel, PhD

Thesis: Tiling 2-deficient Rectangles with Trominoes

Research Experience

Washington University in St. Louis, St. Louis, MO

June 2014 – present

Graduate student, Laboratory of Li Ding, PhD

Thesis: Multi-platform genomics and transcriptomics integration for gene fusion discovery and somatic mutation haplotyping in cancer

- Pan-cancer computational projects utilizing tumor and normal tissue sequencing data to understand patterns of germline and somatic variation
- Integrative data analysis tool and pipeline development, including variant calling and quality control using bulk DNA and RNA, single-cell RNA, and linked-read DNA sequencing
- Gene fusion discovery from The Cancer Genome Atlas (9,624 patients, 33 cancer types) and a large multiple myeloma cohort (742 patients), highlighting clinically-relevant events
- Linked-read whole genome sequencing data analysis for phasing somatic mutations, understanding haplotype structures, and modeling tumor evolution

My work has led to two first author papers published in *Bioinformatics* and *Cell Reports*, one paper under review, and one in preparation. Additionally, I am co-author on four papers published in *Cell, Nature Medicine, Molecular Psychiatry*, and *Annals of Oncology*.

Virginia Commonwealth University, Richmond, VA

August 2013 - May 2014

Biostatistician, Laboratory of Nengliang Yao, PhD

- Led statistical aspects of cancer care research as part of a healthcare policy team
- Procured, cleaned, and analyzed data from national cancer databases, including SEER
- Developed geographically weighted regression and spatial panel data models

This work led to a co-author publication in *PLOS One*.

University of Washington, Seattle, WA

September 2011 – August 2013

Graduate research assistant, Laboratory of Brian Browning, PhD, and Sharon Browning, PhD Thesis: Rare variant method for identity-by-descent detection in sequence data, and the time to most recent common ancestor given identity-by-descent segment length

- Implemented novel algorithms to define identity-by-descent (IBD) between individuals using rare DNA variation and developed statistical models for shared ancestor inference
- Worked with data from the 1000 Genomes Project and simulated coalescent model data
- Assisted in methods development for missing data imputation and haplotype phasing

Presbyterian College, Clinton, SC

Spring 2010

Undergraduate honors research in mathematics, advised by Doug Daniel, PhD Thesis: Tiling 2-deficient Rectangles with Trominoes

• Developed analytical and computational approaches to answer an open question about tiling deficient rectangular fields with trominos

Medical University of South Carolina, Charleston, SC

Summer 2009

Undergraduate Summer Research Fellow, Laboratory of Jim Zheng, PhD

• Created a weighted clustering coefficient model to detect regions of high interconnectivity in yeast gene networks based on synthetic lethal interactions

Presbyterian College, Clinton, SC

Fall 2008 – Spring 2009

Undergraduate research assistant, Laboratory of James Wanliss, PhD

• Analyzed space weather data to refine solar storm prediction models

RESEARCH FUNDING

Paula C. and Rodger O. Riney Blood Cancer Research Initiative Fund

Washington University School of Medicine

Funding since Fall 2017: \$25 million

Principal Investigators include: Li Ding, PhD

My contributions:

- Multiple myeloma gene fusion discovery project, single cell RNA analysis, and linked-read whole genome sequencing analysis
- Preparation and delivery of multiple donor presentations

PUBLICATIONS

- Sanchez-Vega, F., M. Mina, J. Armenia, W. K. Chatila, A. Luna, K. C. La, S. Dimitriadoy, D. L. Liu, H. S. Kantheti, S. Saghafinia, D. Chakravarty, F. Daian, Q. Gao, M. H. Bailey, W. W. Liang, S. M. Foltz, I. Shmulevich, L. Ding, Z. Heins, A. Ochoa, B. Gross, J. Gao, H. Zhang, R. Kundra, C. Kandoth, I. Bahceci, L. Dervishi, U. Dogrusoz, W. Zhou, H. Shen, P. W. Laird, G. P. Way, C. S. Greene, H. Liang, Y. Xiao, C. Wang, A. Iavarone, A. H. Berger, T. G. Bivona, A. J. Lazar, G. D. Hammer, T. Giordano, L. N. Kwong, G. McArthur, C. Huang, A. D. Tward, M. J. Frederick, F. McCormick, M. Meyerson, Cancer Genome Atlas Research Network, E. M. Van Allen, A. D. Cherniack, G. Ciriello, C. Sander and N. Schultz (2018). Oncogenic Signaling Pathways in The Cancer Genome Atlas. Cell 173(2): 321-337 e310
- 2. Gao, Q.*, W. W. Liang*, S. M. Foltz*, G. Mutharasu, R. G. Jayasinghe, S. Cao, W. W. Liao, S. M. Reynolds, M. A. Wyczalkowski, L. Yao, L. Yu, S. Q. Sun, Fusion Analysis Working Group, Cancer Genome Atlas Research Network, K. Chen, A. J. Lazar, R. C. Fields, M. C. Wendl, B. A. Van Tine, R. Vij, F. Chen, M. Nykter, I. Shmulevich and L. Ding (2018). Driver Fusions and Their Implications in the Development and Treatment of Human Cancers. Cell Rep 23(1): 227-238 e223. *Authors contributed equally.
- 3. Foltz, S. M., W. W. Liang, M. Xie and L. Ding (2017). MIRMMR: binary classification of microsatellite instability using methylation and mutations. *Bioinformatics* **33**(23): 3799-3801
- Dang, H. X., B. S. White, S. M. Foltz, C. A. Miller, J. Luo, R. C. Fields and C. A. Maher (2017). ClonEvol: clonal ordering and visualization in cancer sequencing. *Ann Oncol* 28(12): 3076-3082
- Ye, K., J. Wang, R. Jayasinghe, E. W. Lameijer, J. F. McMichael, J. Ning, M. D. McLellan, M. Xie, S. Cao, V. Yellapantula, K. L. Huang, A. Scott, S. Foltz, B. Niu, K. J. Johnson, M. Moed, P. E. Slagboom, F. Chen, M. C. Wendl and L. Ding (2016). Systematic discovery of complex insertions and deletions in human cancers. *Nat Med* 22(1): 97-104
- 6. Olfson, E., N. L. Saccone, E. O. Johnson, L. S. Chen, R. Culverhouse, K. Doheny, S. M. Foltz, L. Fox, S. M. Gogarten, S. Hartz, K. Hetrick, C. C. Laurie, B. Marosy, N. Amin, D. Arnett, R. G. Barr, T. M. Bartz, S. Bertelsen, I. B. Borecki, M. R. Brown, D. I. Chasman, C. M. van Duijn, M. F. Feitosa, E. R. Fox, N. Franceschini, O. H. Franco, M. L. Grove, X. Guo, A. Hofman, S. L. Kardia, A. C. Morrison, S. K. Musani, B. M. Psaty, D. C. Rao, A. P. Reiner, K. Rice, P. M. Ridker, L. M. Rose, U. M. Schick, K. Schwander, A. G. Uitterlinden, D. Vojinovic, J. C. Wang, E. B. Ware, G. Wilson, J. Yao, W. Zhao, N. Breslau, D. Hatsukami, J. A. Stitzel, J. Rice, A. Goate and L. J. Bierut (2016). Rare, low frequency and common coding variants in CHRNA5 and their contribution to nicotine dependence in European and African Americans. Mol Psychiatry 21(5): 601-607

7. Yao, N., S. M. Foltz, A. Y. Odisho and D. C. Wheeler (2015). Geographic analysis of urologist density and prostate cancer mortality in the United States. *PLOS One* **10**(6): e0131578

Honors and Awards

- "Best of *Cell Reports* 2018" co-first author of a frequently cited publication (Driver Fusions and Their Implications in the Development and Treatment of Human Cancers).
- "Outstanding Mentor" recognizing one mentor working with St. Louis high schoolers through the Washington University Young Scientist Program (25th Anniversary Gala, October 22, 2016).
- "Outstanding Senior in Mathematics" awarded to the top graduating senior in the department (Presbyterian College Honors Convocation, April 15, 2010).

TEACHING EXPERIENCE

Washington University in St. Louis, St. Louis, MO

Instructor, Introduction to Python – Bioinformatics

Winter 2019

Student profile: 6 post-baccalaureate students from the Opportunities in Genomics Research diversity outreach program.

Responsibilities: Developed syllabus, created course content, led weekly classroom lectures and activities, gathered regular student feedback.

Teaching assistant, Genomics (Biol 5488)

Spring 2016

Student profile: 40 first-year graduate students from different biomedical programs and with a wide range of programming experience.

Responsibilities: Taught scientific and computational concepts for weekly programming labs (Python), hosted office hours, evaluated student performance on labs and exams (one of four TAs).

Tutor, Fundamentals of Biostatistics for Graduate Students (Biol 5075) Fall 2015, Fall 2016 Student profile: 30 first-year graduate students mostly lacking computational experience. Responsibilities: Taught background bioinformatics and computer coding skills (Python) for programming lab assignments.

Presbyterian College, Clinton, SC

Tutor, Department of Mathematics

January 2007 – April 2010

Student profile: Individuals or groups of undergraduate students from courses up to Calculus II needing help with problem sets and test preparation.

Responsibilities: Re-taught core mathematics concepts in an applied way to bridge gaps in each student's understanding.

MENTORING EXPERIENCE

Mentoring for Ding Lab rotation graduate students

• Yize Li (Fall 2017)

Current status: joined the lab

• Ruiyang Liu (Fall 2017)

Current status: joined the lab

• Lijun Yao (Fall 2017)

Current status: joined the lab

Mentoring for Ding Lab undergraduate students

Moses Schindler (Summer – Fall 2019)
 Current status: Washington University in St. Louis, Class of 2023

• Jessika Baral (Fall 2018)

Current status: Washington University in St. Louis, Class of 2021

• Guanlan Dong (Fall 2018, Spring 2019)

Graduated Washington University in St. Louis: Spring 2019

Current status: graduate student at Harvard University

• Justin Chen (Summer 2018)

Current status: Columbia University, Class of 2022

• Edwin Qiu (Spring 2018, Fall 2018)

Current status: Washington University in St. Louis, Class of 2020

Mentoring for Washington University Young Scientist Program high schoolers

• Jada Reid (Fall 2014 – Spring 2018)

Graduated Collegiate School of Medicine and Bioscience: Spring 2018

Current status: Wesleyan University, Class of 2022

PRESENTATIONS

Oral presentations

- "RNA-Seq and fusion discovery pipeline for a large multiple myeloma cohort." WashU DBBS Precision Medicine Pathway Retreat. October 2017.
- "RNA and DNA methylation signatures for scoring tumor stemness." WashU DBBS Precision Medicine Pathway Retreat. September 2016.

Poster presentations

- "Fusion gene detection across a large cohort of multiple myeloma patients." International Myeloma Workshop. Boston, MA. September 2019.
- "Comprehensive Multi-Omics Analysis of Gene Fusions in a Large Multiple Myeloma Cohort." American Society of Hematology. San Diego, CA. December 2018.
- "Multiple Myeloma RNA-seq and Fusion Discovery Pipeline." WashU DBBS Computational and Systems Biology / Human and Statistical Genetics / Molecular Genetics and Genomics Joint Program Retreat. Potosi, MO. September 2017.
- "Genomic region and sample selection strategy for variant discovery and association analysis." Genome Informatics. Cold Spring Harbor, NY. October, 2015.
- "Comprehensive Analysis of Germline Complex Insertions and Deletions." WashU DBBS Computational and Systems Biology / Molecular Genetics and Genomics Joint Program Retreat. New Haven, MO. September, 2015.
- "Comprehensive Analysis of Germline Complex Insertions and Deletions." WashU DBBS Human and Statistical Genetics Program Retreat. St. Louis, MO. September, 2015.
- "Identity-by-descent analysis of sequence data." International Genetic Epidemiology Society. Stevenson, WA. October 2012.

RESEARCH SKILLS

- Computational project management and data analysis team leadership
- Multi-institutional collaborations, e.g. The Cancer Genome Atlas (TCGA), Clinical Proteomic Tumor Analysis Consortium (CPTAC), Multiple Myeloma Research Foundation (MMRF)
- Analysis of large-scale sequencing data (DNA, RNA) and file formats (VCF, BAM, MAF, BED) on large Unix clusters
- Cancer bioinformatics tools: fusion detection, copy number detection, gene expression, clonal evolution modeling
- Programming languages: R, Python, Bash, Unix command line (github.com/envest)
- Appropriate use of statistical modeling, including association testing and survival analysis
- Data presentation tools: ggplot2 (R package), Adobe Illustrator, LATEX

Professional Development

• Teaching Center

- CIRTL certification
- SEPAL Active Learning seminar
- Mentor training
- Early Career Workshop
- Science Communication class
- CIRTL National Forum

SERVICE AND OUTREACH

Young Scientist Program, Washington University in St. Louis, St. Louis, MO

Leader of Mentoring Program Cohort

Fall 2014 – Spring 2018

- Led 12 graduate students paired with high schoolers for four years
- Coordinated bi-weekly school visits and educational field trips
- Facilitated lesson planning and resources available for hands on science activities
- Mentored one student (Jada Reid, Wesleyan University, Class of 2022)

St. Louis Science Center, St. Louis, MO

Exhibit interpreter Summer 2015

- Smithsonian and NHGRI sponsored exhibit Genome: Unlocking Life's Code
- Discussed the science, ethics, and societal impacts of genomics with museum patrons

Graduate School Coursework

Washington University in St. Louis, St. Louis, MO

June 2014 – present

Ph.D. Biology and Biomedical Sciences (Human and Statistical Genetics) (GPA: 4.00/4.00)
Human Linkage and Association Analysis; Fundamentals of Mammalian Genetics; Computational Statistical Genetics; Genomics; Human Genetics Journal Club; Genetics and Genomics of Disease; Ethics and Research Science; Communicating Science: Writing for Multiple Audiences; Scholarship of Teaching and Learning

University of Washington, Seattle, WA

September 2011 – August 2013

MS Biostatistics (GPA: 3.43/4.00)

Biostatistics I, II; Statistical Inference I, II; Theory of Linear Models; Statistical Genetics I; Categorical Data Analysis; Survival Data Analysis; Evolutionary Genetics; Population Genetics; Statistical Genetics Seminar; Biostatistics Seminar

OTHER WORK EXPERIENCE

Across, Nairobi, Kenya

September 2010 – July 2011

Institutional and Organizational Development Volunteer

Promoted the organization and its mission (holistic transformation of South Sudan communities) to an international audience through the website and other publications. Key aspects of the organization's mission included: serving refugees and internally displaced persons, promoting gender equity and sustainable agriculture, and training teachers and health care providers. Placement coordinated through the Presbyterian Church (U.S.A.) Young Adult Volunteers program.

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